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Vestibular contribution to motor output is also suppressed by voluntary action of the arm

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Abstract figure legend During *Rest, Voluntary reaching* and *Isometric contractions* (Task), participants were subjected to Galvanic Vestibular Stimulation (GVS), Sham and No Stimulation (Stim type), while electromyographic (EMG) activity was recorded from neck and upper extremity muscles (Variables). We estimated intermuscular coherence (Analisis) between muscle pairs to determine the vestibular contribution to arm movements and if vestibular suppression occurs during voluntary arm movements in humans (Outcomes).

Abstract The vestibular sensory system is among the oldest and most fundamental contributors to motor behaviour as it is critical to maintaining posture and balance. However such low-level motor responses could interfere with cortically mediated voluntary behaviour that naturally involves posture and balance. Consequently, it has been proposed that - much like the inhibition of reflex responses - vestibular contributions to motor output are 'gated' (dubbed vestibular suppression) to avoid undesirable self-perturbations during voluntary head movements. Here we demonstrate that such suppression also occurs for unperturbed voluntary arm function. Our evidence comes from comparing coherence at baseline (No Stimulation) and after Sham and Galvanic Vestibular Stimulation (GVS). Specifically neck muscles showed shared neural drive (intermuscular coherence [IMC]), which increased with GVS - but not Sham - at Rest. This GVS-mediated increased coherence in neck muscles, however, was suppressed during voluntary isometric contractions and reaching movements of the arm on the same side as the GVS was applied. No changes were found in pairwise IMC during Sham (compared with No stimulation) or in arm muscles either at rest or during voluntary movement during GVS in neurotypical adults. In addition to extending vestibular suppression to unperturbed voluntary arm function, these results provide support for the common (yet unproven to our knowledge) notion that arm muscles do not receive vestibular neural drive during unperturbed voluntary movement. Moreover, these results shed light on the mechanisms that mediate competing descending outputs for voluntary function and serve as a baseline against which to compare potential task-dependent dysregulation of vestibular-mediated output to the neck and arms in stroke and neurological conditions.

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Key points

- The vestibular system is critical for correcting perturbations during voluntary movement.
- During voluntary head movements, vestibular suppression occurs to avoid undesirable self-perturbations.
- However, the contribution of the vestibular system to unperturbed voluntary arm movement remains unclear.
- We used intermuscular coherence (IMC) to measure vestibulospinal drive to neck and arm muscles while applying Galvanic Vestibular Stimulation (GVS), Sham and No Stimulation. We compared IMC at Rest and unperturbed voluntary movement of the arm in neurotypical adults.
- Neck muscles showed increased shared neural drive at rest, only when GVS was applied. However, vestibular drive was suppressed during unperturbed voluntary isometric contractions and reaching movements of the arm.
- Vestibular drive to arm muscles did not increase when GVS was applied.
- We provide evidence that arm muscles do not receive vestibulospinal drive, excluding its contribution to unperturbed voluntary movement.
- These results could provide valuable insights into the vestibular contribution to motor impairments following neurological conditions such as stroke.

Introduction

The otoliths and semi-circular canals are constantly sensing expected and unexpected head orientation and movement relative to space (Cullen, 2023a, 2023b). These inputs are integrated with somatosensory and predictive self-motion signals from the brainstem, cerebellum and cortex (Cullen & Zobeiri, 2021). Such integration is essential for reflexive stabilization of gaze and posture, and accurate control of voluntary movements (Cullen, 2023a, 2023b).

While vestibular reflexes are essential for providing robust responses to unexpected external stimuli, they could be counter-productive when they interfere or compete with motor signals for voluntary movements (Lopez & Cullen, 2024; Niyo et al., 2024). Consider how vestibular reflexes can be in competition with voluntary function when, for example, voluntary head rotations need to be ignored during reaching movements. Experimental evidence shows that these reflexes - observed as electromyographic responses to perturbations - are largely suppressed during active head movements (Cullen, 2023a, 2023b; Kwan et al., 2019). Notably, responses in the vestibular nuclei are suppressed (vestibular suppression) and occur in neck muscles during voluntary and self-initiated head movements (Cullen & Zobeiri, 2021). Importantly, vestibular suppression in leg muscles is also seen and modulated during voluntary leg movements for locomotion (Dakin et al., 2013) - presumably to avoid motor interference. On the contrary the contribution of vestibular drive to upper extremity voluntary movement is critical for sensing our self-initiated movements relative to the environment (Cullen, 2023a). Moreover, it allows us to estimate additional physical forces (e.g. coriolis and centrifugal forces) needed to plan and execute an accurate movement, such as during reaching. Accordingly, the motor pathways controlling reaching movements demonstrate feedback-mediated responses at a minimal latency of 50 ms. These responses compensate for displacements of the body and limb - relative to a reaching target produced by externally applied perturbations as well as during self-motion (Adamovich et al., 2001; Azadjou

et al., 2023). Moreover, they are reduced after unilateral vestibular lesions, excluding their emergence from the proprioceptive system (Raptis et al., 2007). In fact, the vestibular system contributes to the high-level planning of reaching movements, which is crucial for achieving accurate movement performance (Azadjou et al., 2023; Klam & Graf, 2006; Schlack et al., 2002). For example, neurons in the macaque parietal cortex show increased firing responses to vestibular signals, which are integrated with other somatosensory inputs, including proprioception, vision and touch (Cullen, 2023b). Furthermore, the corticospinal tract and brainstem output - from the medial longitudinal fasciculus (MLF) which contains reticulo-, vestibulo- and tecto-spinal tracts - send converging and overlapping signals to the spinal cord during reaching and grasping tasks (Riddle & Baker, 2010).

The contribution of the vestibular system to voluntary movement is determined by measuring vestibulospinal drive while applying GVS. Vestibular afferents are stimulated through current applied via transmastoid surface electrodes, increasing vestibulospinal drive, without affecting proprioception and tactile sensory information (Cullen, 2023b; Forbes et al., 2015; Kwan et al., 2019). The GVS-mediated increase in vestibular drive evokes both ocular and postural responses with electromyographic responses in axial and appendicular muscles with a latency of 8–50 ms (Cullen, 2023a; Forbes et al., 2015).

Given that upper limb movement also depends on accurate suppression of reflexes and accurate estimation of position and velocity of the body, we investigated the vestibular contribution to arm movements and whether suppression is also a mechanism to enable voluntary arm movements in humans. We hypothesize that, as in voluntary neck and leg movements, vestibular contributions to the activation of neck and arm muscles should differ between rest and voluntary movement of the arm. Understanding the role of the brainstem vestibular output in arm movements could then provide valuable insights into its contribution to motor impairments following neurological conditions such as stroke.

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Ethical approval

The study conformed to the standards set by the Declaration of Helsinki, except for registration in a database. All participants gave their informed written consent to participate in this study, which was approved by the University of Southern California Internal Review Board (USC IRB: HS-17-00304).

Study participants

Seventeen right-handed individuals participated in the study (n = 17; 7 males; 10 females), with a mean age of 21.5 years (ranging from 18 to 27 years), all free from pain, injury or any conditions affecting upper limb movement. Importantly, all participants were free from any neurological condition affecting control of the upper extremity (neurotypical).

Tasks

Participants performed the following tasks while sitting. *Rest*: Participants were seated with their hands resting on their lap or armrest. They were encouraged to stay relaxed and silent for 90 s at the beginning of the experimental procedures to collect baseline muscle activity (Fig. 1, left panel). *Voluntary reaching*: Participants were seated in front of a hand-powered ergometer mounted to be rotated in the horizontal plane with their right arm (Fig. 1, middle panel). The protocol for this task is thoroughly described in a previous article (Laine et al., 2021).

Isometric contraction: Participants were seated with the backrest inclined 15° backwards (Fig. 1, right panel). They

were encouraged to keep their heads as close as possible to the headrest, without supporting any weight on it. This position was enforced to have a bilateral isometric contraction of the sternocleidomastoid muscles (SCMs) and Upper Trapezius (UTrap) for head stabilization. Simultaneously, they held a 2.26-kg dumbbell with their hands, while their forearm was kept parallel to the ground and the arm parallel to the inclined backrest, but without any support on it. This position induced an isometric contraction of the Biceps Brachii, Anterior and Middle Deltoid muscles. Participants were encouraged to hold this position for 90 s, while verbal feedback was provided to correct or return to the instructed position if they departed from it. They were allowed to rest or to support their heads/extremities if they felt fatigued; however, it was

Stimulus types

not needed by any participant (Fig. 2).

For each task, participants were subjected to three stimulus types as follows: No stimulation, GVS and Sham stimulation. Based on previous protocols on human participants, GVS consisted of a binaural galvanic stimulation where the positive electrode was placed on the right mastoid process (negative electrode on the left mastoid process) (Forbes et al., 2015). The position of the electrodes was chosen to induce a vestibular response on the right side of the body, which was confirmed by visual inspection of the EMG signal from the SCM muscle (see Fig. 3, lower left panel). The stimulation frequency was set at 4 Hz, with an amplitude ranging between 0.8 and 1.2 mA. The amplitude was modulated to avoid EMG signal saturation from the SCM muscle



Figure 1. Tasks performed by each participant

Left: At the beginning of the experiment, each participant sat comfortably in a chair with their hands on their lap, while EMG was recorded at *Rest*. Middle: During the *Reaching* task, participants were asked to rotate a horizontal ergometer in a counterclockwise rotation with their right arm to produce a cyclical movement. Right: In the *Isometric* condition, subjects were positioned with a 15° inclined backrest, while raising their head from the headrest to activate the SCMs muscles. Meanwhile, their right upper extremity remained unsupported as they held a 2.26-kg dumbbell, with their forearm parallel to the ground and their arm aligned with the backrest.

or if the participant felt uncomfortable. Independent of the stimulation amplitude, the Sternocleidomastoid response to GVS was always clearly visible and greater than the EMG signal at rest. For the Sham stimulation, a mechanical vibration (400 Hz) was delivered on the right mastoid (same location as the positive GVS's electrode). Each participant completed nine randomized conditions: three tasks (Reaching, Isometric and Rest) and three stimulus types (GVS, None and Sham). To mitigate potential carryover effects of GVS and Sham, the Rest + No stimulation condition was always completed first. Moreover, to assess the potential carryover effects of GVS, the resting condition was repeated at the end of the experiment. Subsequently, the Rest + No stimulation conditions at the beginning and end of the experimental procedures were compared, to determine if the vestibular drive remains increased over time, even after GVS has ceased.

Data acquisition and processing

A custom game was designed in c# to collect the angle data from the ergometer and provide live real-time feed-

back about the user's rotation velocity (Unity3D, San Francisco, CA, USA). Custom hardware provided a pulse via an Arduino MEGA (Arduino, Somerville, MA, USA) to synchronize EMG data, angle measurements and GVS stimuli delivery time. We collected EMG signals at 2.5 kHz from seven muscles of the right upper extremity using a DataLINK system and associated software (Biometrics Ltd, Newport, UK). Surface EMG sensors (Biometrics Ltd SX230: bipolar, gain: 1,000, bandwidth: 20-460 Hz) were placed over the right arm: Biceps Brachii (Bic), lateral head of the Triceps Brachii (Tric), Anterior, Middle and Posterior Deltoid (ADelt, MDelt, PDelt, respectively), UTrap and SCMs, following standard recommendations from SENIAM. Electrode placement and signal quality were confirmed using palpation of each muscle and observation of the EMG during voluntary activation. This set of muscles is sufficient for a general analysis of coupling among the shoulder/elbow muscles relevant to our task (Laine et al., 2021). All EMG signals were processed offline using R/Rstudio (R Core Team, 2021).

To remove GVS artefacts, a 10-ms window surrounding each stimulus (from 2 ms before to 8 ms after the electrical pulse) was replaced with empty or missing values from



Figure 2. Raw and Processed (full-wave rectification and band-pass filtered) EMG signals from a typical participant without stimulation, recorded during the *Isometric* task, 5 s after the task started Each trace represents 2 s of data. The left panel displays the *Raw* and *Processed* EMG signals for the *Sternocleidomastoid* (SCM) and *Upper Trapezius* (UTrap) muscles. The right panel overlays the signals with the sequential processing steps: the light trace corresponds to the Raw signal (see positive and negative values), and the darkest trace represents the final processed signal.

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the SCM, Deltoid (ADelt, MDelt, PDelt) and UTrap EMG signals (see Fig. 3, bottom panel). The missing data points were then interpolated to prevent aliasing and preserve the signal's frequency characteristics. Because vestibular responses have a latency of 8-50 ms, we prevented their removal from the signals during the replacing and interpolation process (Forbes et al., 2015). Signal processing was performed according to our paper published previously (for a detailed description, see Laine et al. 2021). In summary, all EMG signals were downsampled to 1000 Hz, band-pass filtered between 8 and 250 Hz and then full wave rectified. The filter's purpose was to remove any remaining artefacts arising from GVS, as well as from those frequencies irrelevant for intermuscular analysis. EMG rectification was done to enhance intermuscular coherence (IMC) and avoid distortion of motor synchronization (Boonstra & Breakspear, 2012).

Statistical analysis

Coherence analysis measures the shared neural drive between two signals in the frequency domain (correlation in the frequency domain). As such, IMC assesses the degree of synchronization between the neural drive to two muscles on the basis of their EMG signals (Boonstra, 2013). Consequently, we calculated magnitude squared coherence between each muscle pair using 300-ms windows and a 50% overlap (Laine et al., 2021).

A threshold to determine significant pairwise coherence was built for easy visual inspection (see red dotted line in Figs. 4 and 5). To estimate the coherence expected by chance, we generated 1000 phase-randomized surrogate series for each muscle pair and participant following the methods described by Ebisuzaki (1997). Coherence was calculated for each surrogate pair. A 95% confidence interval (95% CI) was constructed for



Figure 3. The *Sternocleidomastoid* (SCM) muscle shows a stereotypical response across all tasks, whereas the *Upper Trapezius* (UTrap) shows a smaller response restricted to *Rest*, emphasizing its suppression during *Isometric* and *Reaching* tasks

Top: Raw and Processed (GVS artefact removed, full-wave rectified and band-pass filtered) EMG activity from *Sternocleidomastoid* (SCM) and *Upper Trapezius* (UTrap) muscles during each task (when *GVS* was applied) for a typical participant. Each trace corresponds to one second of activity, which was extracted when the signal was stable (5 s after the tasks started) *Bottom*: Mean (\pm SD) Raw and Processed EMG activity (across stimuli) in response to Galvanic Vestibular Stimulation (*GVS*), from 50 ms before to 110 ms after the stimuli was delivered. For visualization purposes of the stereotypical response, all stimuli were aligned to their delivery time (t = 0). It showcases the averaged EMG responses across all stimuli from a single participant (number of averaged stimuli highlighted inside each plot). The light trace shows the GVS artefact, whereas the darker trace shows the EMG after this artefact was removed and the signal processed (full-wave rectified and band-pass filtered). The shaded grey area represents the 10-ms window (ranging from -2 to 8 ms) during which the GVS artefact was removed before signal processing (see 'Materials and methods' section). Following this window, suppression of the UTrap response is observed during voluntary action of the arm, as opposed to SCM responses, which are always visible.

each muscle pair based on the *z*-transformed coherence values obtained from the surrogates. This interval was built across all frequency bands to provide a stringent and robust criterion for assessing statistical significance during visual inspection. Importantly, this 95% CI was not used in hypothesis testing but rather as a visual reference to highlight when individual pairwise coherence exceeds the highest values expected by chance. In other words, individual pairwise coherence above the 95% CI is likely to be significant, revealing true shared drive between the two muscles.

For unbiased statistical estimation during hypotheses testing, all raw – magnitude-squared – coherence values were transformed to *z*-scores (IMC_z) using Fisher's Z transform (IMC_z = atanh(rho)) before performing statistical comparisons (Laine et al., 2021). We compared IMC_z across the three stimulus types and tasks. The values of IMC_z across the frequency spectrum were gathered into three bands as follows: alpha (8–16 Hz), beta (16–30 Hz) and gamma (30–50 Hz) frequency bands. This approach allowed us to minimize statistical error by decreasing the number of *post hoc* comparisons to make if the null hypothesis was rejected. Given the lack of a non-parametric alternative to test our hypotheses, a Robust Mixed Effect ANOVA model (i.e. robust repeated Measures ANOVA model) was used to test if GVS increased neural drive. The assumptions of the classical ANOVA statistical tests are not required to be met when performing these robust statistical methods. Then, if a GVS effect was found across frequency bands,



Figure 4. During Rest, Galvanic Vestibular Stimulation (GVS) increases intermuscular coherence (IMC) across alpha, beta and gamma frequencies between neck muscles, but not neck muscles We compared *z-transformed* IMC coherence across frequency bands during Rest under three different stimulus types: No Stimulation (*None*), *Sham* and GVS. Neck muscles are *Sternocleidomastoid* (SCM) and *Upper Trapezius* (UTrap). Arm muscles are *Biceps brachii* (Bic) and *Anterior Deltoid* (ADelt). Red dotted line indicates the 95% upper confidence interval estimated from 1000 randomizations of the original signals. The middle and lower panels include the mean IMC_z across stimulus types on top of their corresponding box for easy comparison while keeping the same scales across muscle pairs.

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we compared IMC_z using statistical parametrical mapping (SPM) across the three stimulus types for each task separately (*Rest, Reaching* and *Isometric*) to identify the specific frequencies at which the differences were significant. Finally, to test for a carryover effect (i.e. long-lasting GVS effect), we compared coherence at the beginning and end of the experimental protocol when the participant was at rest and without any stimulus. Post hoc comparisons were made using Dunnett test with Bonferroni corrections (adjusted for two comparisons: GVS vs. No Stimulation and GVS vs. Sham) (Table 1).

Results

As expected, the shared neural drive between neck muscles at *Rest* (as per IMC_z) increased during *GVS* (Fig. 4). This was confirmed by a repeated measures ANOVA model with *post hoc* comparisons and Bonferroni corrections at each frequency band (i.e. alpha to gamma, see Fig. 4 and Table 2). Importantly, no significant differences were found in IMC_z for neck muscles between None and Sham stimulus types (*P*-values: alpha = 0.76, beta = 0.7, gamma = 0.55; see Table 2). This supports the notion that GVS increases vestibular output only to neck



Figure 5. Galvanic vestibular stimulation (GVS, blue line) increases vestibular drive to neck muscles (SCM-UTrap) at Rest

However, it is suppressed during Reaching and Isometric tasks. GVS did not increase vestibular output to neck-arm and arm muscles (SCM-Bic and Bic-MDelt, respectively), neither at Rest nor during unperturbed voluntary action of the arm. No differences were found between Sham and No stimulation (which excludes a placebo effect). Mean (\pm SD) *z*-transformed IMC_z coherence across subjects during rest condition under three different stimuli: No Stimulation (*None*), *Sham* and Galvanic Vestibular Stimulation (*GVS*). Each plot (except for SCM-UTrap at *Rest*) includes an inset where the signals are auto-scaled. Neck muscles are SCM: *Sternocleidomastoid* and UTrap: *Upper Trapezius*. Arm muscles are Bic: *Biceps Brachii* and MDelt: *Middle Head of Deltoid*. Values above the dotted red line can be considered to have significant IMC_z, as per the 95% upper confidence interval estimated from 1000 randomizations of the original signals. Alpha, beta and gamma frequency bands are identified by blue vertical dashed line on top of upper panel.

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	Alpha			Beta			Gamma		
Muscle	<i>F</i> -value (DF ₁ , DF ₂)	P-value	η^2	F-value (DF ₁ , DF ₂)	<i>P</i> -value	η^2	<i>F</i> -value (DF ₁ , DF ₂)	P-value	η^2
SCM-UTrap (N-N)	7.35 (1.5, 24.6)	0.01	0.31	17.16 (1.2, 19.8)	< 0.001	0.52	9.35 (1.2, 19.3)	< 0.001	0.37
SCM-Bic (N-A)	1.62 (1.6, 25.7)	0.22	0.09	0.79 (1.8, 29.4)	0.45	0.05	0.95 (1.6, 26.3)	0.38	0.06
UTrap-Bic (N-A)	1.04 (1.2, 19.8)	0.34	0.06	1.07 (1.2, 19)	0.33	0.06	1.09 (1.1, 17.9)	0.32	0.06
UTrap-Tric (N-A)	1.34 (1.6, 25.7)	0.27	0.08	0.88 (1.2, 19.5)	0.38	0.05	1.54 (2, 32)	0.23	0.09
UTrap-MDelt (N-A)	0.63 (1, 16.6)	0.45	0.04	0.89 (1, 16.2)	0.36	0.05	0.98 (1, 16.7)	0.34	0.06
Bic-ADelt (A-A)	1 (1.7, 27.1)	0.37	0.06	1.46 (1.3, 21.6)	0.25	0.08	1.43 (1.3, 20.9)	0.25	0.08
Bic-MDelt (A-A)	0.48 (1.4, 23.1)	0.57	0.03	0.47 (1.4, 23.1)	0.57	0.03	0.57 (1.4, 22.1)	0.51	0.03
Bic-PDelt (A-A)	0.08 (1.8, 28.3)	0.90	0.00	0.01 (1.7, 27.5)	0.99	0.00	0.02 (1.5, 24.1)	0.95	0.00
ADelt-MDelt (A-A)	1.09 (1.4, 22.6)	0.33	0.06	0.29 (2, 32)	0.75	0.02	0.66 (2, 32)	0.52	0.04
ADelt-PDelt (A-A)	0.39 (1.2, 19)	0.58	0.02	0.11 (1.6, 25.9)	0.85	0.01	0.28 (1.6, 26)	0.71	0.02
MDelt-PDelt (A-A)	0.1 (1.2, 18.6)	0.79	0.01	0.38 (1.1, 17.9)	0.57	0.02	1.14 (1.2, 19.2)	0.31	0.07

Table 1. Robust repeated measures ANOVAs: degrees of freedom, F ratio, P-value and effect size (η^2)

Muscles: Sternocleidomastoid (SCM); Upper Trapezius (UTrap); Biceps brachii (Bic); Triceps brachii (Tric) and Anterior, Middle and Posterior Deltoid (ADelt, MDelt, PDelt). Subscripts indicate Neck–Neck muscles: N-N, Neck–Arm muscles: N-A and Arm–Arm muscles: A-A.

Table 2 Ad	justed P-values for	nost hos com	naricone ucine	Bonforroni	corrections (a	diusted for two	comparisons)
Table Z. Au	Justeu F-values IOI	post not com	parisons using	j bomenom	corrections (a	ujusteu ioi two	companisons)

	Alpha			Beta			Gamma		
Muscle	None/GVS	None/Sham	GVS/Sham	None/GVS	None/Sham	GVS/Sham	None/GVS	None/Sham	GVS/Sham
SCM-UTrap (N-N)	<0.001	0.76	0.02	<0.001	0.70	<0.001	<0.001	0.55	0.01
SCM-Bic (N-A)	0.12	0.69	0.08	0.22	0.96	0.24	0.22	0.75	0.18
UTrap-Bic (N-A)	0.60	0.31	0.30	0.43	0.34	0.29	0.52	0.32	0.29
UTrap-Tric (N-A)	0.11	0.67	0.17	0.28	0.49	0.28	0.10	0.79	0.19
UTrap-MDelt (N-A)	0.38	0.52	0.15	0.37	0.35	0.85	0.31	0.37	0.37
Bic-ADelt (A-A)	0.25	0.76	0.35	0.09	0.64	0.23	0.18	0.50	0.23
Bic-MDelt (A-A)	0.27	0.80	0.45	0.34	0.72	0.44	0.19	0.79	0.40
Bic-PDelt (A-A)	0.72	0.82	0.86	0.94	0.99	0.91	0.93	0.85	0.86
ADelt-MDelt (A-A)	0.36	0.57	0.25	0.87	0.45	0.57	0.77	0.25	0.45
ADelt-PDelt (A-A)	0.52	0.67	0.56	0.70	0.78	0.80	0.54	0.88	0.61
MDelt-PDelt (A-A)	0.85	0.85	0.22	0.62	0.46	0.64	0.26	0.33	0.63

Statistical results (*p*-values) for differences between stimulus types: No Stimulation (*None*), *Sham* and Galvanic Vestibular Stimulation (*GVS*). Muscles: *Sternocleidomastoid* (SCM), *Upper Trapezius* (UTrap), *Biceps brachii* (Bic), *Triceps brachii* (Tric) and *Anterior, Middle and Posterior Deltoid* (ADelt, MDelt, PDelt). Subscript indicates Neck–Neck muscles: *N-N*, Neck–Arm muscles: *N-A* and Arm–Arm muscles: *A-A*.

muscles (see upper panel in Fig. 4 and upper left panel in Fig. 5).

A more detailed analysis of these GVS-driven increases in IMC_z between SCM-UTrap suggests that such vestibular output spans a broad frequency spectrum. This was quantified by SPM analysis in 0.5-Hz bins across frequencies, which showed consistently increased IMC_z during GVS when compared to None and Sham stimulus types. Specifically, IMC_z was higher in the range from 12 and 50 Hz (high alpha to gamma bands, SPM{F}_{2,32}, Random Field Theory threshold = 6.6, P < 0.001, Fig. 5).

A secondary analysis confirmed that our block-randomized application of the three stimulus types did not produce measurable carry-over effects. This

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was confirmed by comparing the baseline coherence of Neck muscles during *Rest* + *No stimulation* conditions at the beginning and the end of the experimental protocol (P = 0.3).

Interestingly, GVS did not increase IMC_z between neck muscles (SCM-UTrap) during unperturbed voluntary *Reaching* or *Isometric Contraction*. This is supported by the results of both repeated measures ANOVA and SPM analysis, which did not reveal significant differences when comparing GVS with either Sham or No stimulation (see middle and right upper panels in Fig. 5). This is an example of vestibular suppression that, to the best of our knowledge, has not been reported during voluntary function in human or non-human primates. Finally, vestibular output is neither present nor suppressed in arm muscles in any condition (*Rest, Reaching* and *Isometric*), demonstrated by no significant differences in IMC_z from muscle pairs in the active arm. This was supported by the results of both repeated measures ANOVA and SPM analysis (see middle and lower panels in Figs. 4 and 5).

Discussion

Our IMC results – as per IMC_z – indicate that neck, but not arm muscles, receive shared neural drive from the vestibular system (brainstem vestibular output) in neurotypical participants. Importantly, this vestibular output to neck muscles is increased at Rest during GVS (see upper panels in Fig. 4 and left upper panel in Fig. 5), but suppressed during voluntary action of the arm (Reaching and Isometric contraction, see middle and right upper panels in Fig. 5), without reaching statistical significance during voluntary movement of the arm. In support of this central result, we did not see increases in IMC_z during the Sham condition when compared with No stimulation. Consequently, we can exclude a tactile or proprioceptive mechanism for the changes seen during GVS, as well as a placebo effect. We propose that this is a previously unreported form of vestibular suppression due to voluntary action of the arm - which was known to exist in neck muscles during head movements (Cullen & Zobeiri, 2021). Moreover, the lack of increase in IMC_z with GVS between neck and arm muscles (SCM-Bic) and between arm muscles (Bic-MDelt) provides evidence that these arm muscles are unlikely to receive vestibular output at Rest or during unperturbed Reaching and Isometric tasks (see Figs. 4 and 5). We conclude that this previously unknown distribution and task-dependent suppression of brainstem vestibular output in neck versus arm muscles during Rest and voluntary function sets a critical baseline in neurotypical adults. Given the clinically practical nature of GVS and EMG recordings of the arm, this baseline can be used to quantitatively assess disruptions of cortical, vestibular and brainstem output in stroke survivors.

Although there are limitations in our study, we believe that these do not detract from our main results or conclusions. A potential limitation of our approach is the phase-randomization method used to estimate the confidence interval for the coherence expected by chance. Although this method preserves the amplitude spectrum, it may not fully account for the temporal and frequency-dependent physiological characteristics of the original signals. This could result in a null distribution for coherence that does not perfectly capture the complexities of physiological signals. We used this method for visualization purposes only, and thus – even if we had underestimated this threshold – it does not change the interpretation of our results. Another potential limitation is that the suppression of responses during voluntary arm movements could be attributed to a lower signal-to-noise ratio (SNR). However, we believe this is unlikely, as the UTrap response is only visible during Rest, and the EMG signal drops during voluntary arm movement (see Fig. 3). Second, the noise in our signal is relatively low. For example, we observe that during Rest, the baseline activity is minimal when compared with voluntary action (see Fig. 3). Therefore, any activity above the levels of the resting condition must be attributed to signal arising from neural drive rather than noise. This implies that vestibular drive is not clearly identified or was suppressed by neural drive originating from other cortical and subcortical sources involved in voluntary movement. Consequently, we believe that the suppression of vestibular responses during voluntary action is likely due to task-related neural modulation rather than an artefact of SNR. We also acknowledge that our results might be explained as a direct stimulation of neck muscles, or stimulation of the XI (or Accessory) cranial nerve (innervating SCM and UTrap muscles), which might be feasible due to the proximity between the GVS electrode and the neck muscles. However, it has been found that current density decreases to 10% at a distance of 10 mm from stimulating surface electrodes (Enoka et al., 2020), which is a small fraction of the distance between the stimulating electrode (behind the ear) and the SCM's innervation point by the XI cranial nerve, which is 4-9 cm below the mastoid process (Bordoni et al., 2018). Moreover, we would not have seen that any vestibular suppression had such spillover stimulation short-circuited brainstem vestibular output to neck muscles. Lastly, our removal of stimulation artefacts up to 8 ms after each stimulation pulse removes, by design, the effect of such direct stimulation as well (Pinto & De Carvalho, 2008). However, this removal of the stimulation artefact every 250 ms may explain the higher (and potentially artefactual) coherence below 8 Hz in Fig. 5.

Vestibular suppression during voluntary function is limited to neck muscles. Vestibular suppression in neck muscles has been previously described during active head movements across several species, including human and non-human primates (Cullen, 2023b). The purported utility is to suppress potentially counter-productive (involuntary) responses that could act as internal perturbations during voluntary movement (Cullen & Zobeiri, 2021; Cullen, 2023a). It has been proposed that vestibular suppression in neck muscles occurs when the consequences of head movement (informed by efference copy and predicted by the cerebellum) are analogous to the expected sensory information received from proprioceptors (Cullen & Zobeiri, 2021; Cullen, 2023a). Accordingly, we interpret the suppression of coherence between neck muscles (during GVS but also during no stimulation, Fig. 5) as a mechanism to prevent the disruptive effect of vestibular output on head and neck kinematics during voluntary action of the arm. Case in point, our experimental design required participants to visually track a dolphin on a screen to maintain the cadence of the crank during *Reaching*, or maintain neck posture during *Isometric Contraction*. These natural tasks required active control of head and neck kinematics which could be perturbed by vestibular responses.

Vestibular output to arm muscles is absent across tasks.

Vestibular output is known to affect the involuntary or reactive activation of arm muscles during perturbations of the trunk (Adamovich et al., 2001). However, our findings suggest that such vestibular-mediated adjustments are not involved or necessary during the self-initiated voluntary tasks we tested which provided stable and static trunk support. Acoustic startle studies suggest that stroke survivors exhibit greater responses on their more-affected (i.e. spastic-paretic) arm, presumably due to increased vestibular output (Miller & Rymer, 2017). That is, the evoked latencies were shorter and strongly lateralized in stroke survivors, which was interpreted as being triggered from vestibular instead of acoustic pathways (Miller & Rymer, 2017). Presumably, these increased vestibular-mediated responses result from the disruption of the inhibitory cortical pathways to brainstem and spine, such as those mediating vestibular projections (Miller et al., 2014; Miller & Rymer, 2017). This lack of vestibular output to arm muscles in our neurotypical participants serves as a baseline for future studies to understand the disruption of pathways affecting arm control in hemiparetic stroke.

Our results support prior reports that IMC of vestibular origin is broadly distributed across the frequency spectrum. IMC in the beta band has been proposed as a biomarker that primarily reflects CST integrity (Fisher et al., 2012; Fang et al., 2009; Ko et al., 2023; Larsen et al., 2017; Mima et al., 2001). On the contrary, the alpha band is thought to do the same for RST and propriospinal pathways (Grosse & Brown, 2003; Thevathasan et al., 2012; Tattersall et al., 2014). Our results during GVS, however, do not show a particular dominance of alpha, beta or gamma frequency bands. This supports prior work making the same inference. For example, during postural tasks, the SCM muscle has been shown to respond to stimulation with increased IMC in frequencies spanning up to 70 Hz, while lower limb muscles were coherent under 25 Hz (Forbes et al., 2015). Blouin et al. (2011) showed that during GVS, ankle muscles - such as gastrocnemius and soleus -

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are coherent during the stance phase of locomotion at frequencies of around 20 Hz. Moreover, during physical activity such as running and jumping, the frequency range of vestibular-mediated responses can increase to 70 Hz (from alpha to gamma bands) (Carriot et al., 2014; Forbes et al., 2015). In agreement with these previous findings, we found increased coherence from alpha to gamma frequency bands at Rest during GVS. Importantly, neural pathways might be reflected and overlapping across these frequency bands. For example, VST and CST drive are both reflected on beta band, whereas RST and VST are both reflected on the alpha band. Therefore, changes in coherence on any given frequency should be interpreted carefully, especially under neurological conditions. This underscores the need to carefully interpret IMC_z in specific frequencies, particularly in stroke. That is, if the disruptions in descending pathways to the more affected arm have contributions from vestibular output, these changes in IMC_z may be reflected across a broad spectrum of frequency bands, masking decreased drive from such disrupted pathways.

In summary, our study demonstrated an increased neural drive to neck, but not arm muscles as a result of vestibular stimulation. The increased vestibular drive at rest was suppressed during isometric contraction and unperturbed voluntary reaching. These findings suggest that vestibular suppression - which has been described in head and neck muscles - also applies to voluntary arm movements. Furthermore, the absence of changes in neural drive to arm muscles during GVS provides evidence to exclude a vestibular contribution to unperturbed voluntary arm movement. Our results establish a baseline for the expected neural drive to arm and neck muscles, which could be valuable in clinical practice for quantitatively assessing disruptions of cortical, vestibular and brainstem output in stroke survivors and people with neurological conditions.

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Additional information

Data availability statement

The data that support the findings of this study are available from the corresponding author upon reasonable request.

Competing interests

The authors declare that this research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Author contributions

A.B.-J. and F.J.V.-C contributed to conception and design of the study. A.B.-J. performed the experiments, pre-processed the data and performed the statistical analysis. A.B.-J. and F.J.V.-C interpreted the results and wrote the manuscript. Both authors contributed to manuscript revision, read and approved the submitted version. All authors have approved the final version of the manuscript and agreed to be accountable for all aspects of the work. All persons designated as authors qualify for authorship, and all those who qualify for authorship are listed.

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Keywords

Galvanic Vestibular Stimulation, vestibular output, voluntary reaching

Supporting information

Additional supporting information can be found online in the Supporting Information section at the end of the HTML view of the article. Supporting information files available:

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